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Truncating Mutations in UBAP1
Cause Hereditary Spastic Paraplegia

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The diagnostic gap for rare neurodegenerative diseases is still considerable, despite continuous advances in gene identification. Many novel Mendelian genes have only been identified in a few families worldwide. Here we report the identification of an autosomal-dominant gene for hereditary spastic paraplegia (HSP) in 10 families that are of diverse geographic origin and whose affected members all carry unique truncating changes in a circumscribed region of UBAP1 (ubiquitin-associated protein 1). HSP is a neurodegenerative disease characterized by progressive lower-limb spasticity and weakness, as well as frequent bladder dysfunction. At least 40% of affected persons are currently undiagnosed after exome sequencing. We identified pathological truncating variants in UBAP1 characterized by progressive lower-limb spasticity and weakness, as well as frequent bladder dysfunction. At least 40% of affected persons are still not diagnosed even after whole-exome sequencing (WES). Furthermore, many of the genes reported in recent years have only been described in a few families.1

In an effort to further close this diagnostic gap in HSP, we have gathered a highly diverse sample of 10 families from six countries (Iran, 1; USA, 1; Germany, 4; Canada, 1; Bulgaria, 2; and Spain, 1). Prior to the initiation of this study, all participating affected individuals gave informed consent in agreement with each institutional review board. In one family of Persian origin, family 1, we were able to genetically ascertain a total of 14 affected individuals from three generations (Figure 1). Sequencing of an HSP gene panel and CNV

Hereditary spastic paraplegia (HSP) represents a group of genetically highly heterogeneous rare inherited neurodegenerative diseases, which are characterized by the pathological hallmark of a length-dependent degeneration of corticospinal-tract axons (see GeneReviews in Web Resources).1 Clinically, HSPs are marked by progressive spastic paraparesis, although the clinical presentation encompasses a wide spectrum of phenotypes. In pure forms of HSP, progressive spasticity and weakness in the lower extremities are the main features. In complex forms of HSP, additional clinical symptoms include cataracts, ataxia, epilepsy, cognitive impairment, peripheral neuropathy, optic neuropathy, and deafness (see GeneReviews in Web Resources).1 The prevalence of HSP has been estimated to be 1.3–9.6 in 100,000 (see GeneReviews in Web Resources).1,2 Thus far, at least 58 genes have been reported to cause HSP in a Mendelian fashion.3 Yet approximately 40% of affected persons are still not diagnosed even after whole-exome sequencing (WES). Furthermore, many of the genes reported in recent years have only been described in a few families.1

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analysis at the SPG4 locus were unremarkable. Subsequently, WES was performed in two affected individuals (V.1 and V.15). Bioinformatics analysis of the sequencing data used standard tools, including BWA aligner,\(^5\) FreeBayes,\(^5\) GATK,\(^6\) and GENESIS.\(^7\) Only non-synonymous variants with a minor-allele frequency of less than 0.0001 in gnomAD and in our in-house Iranian variant database (BayanGene; http://www.bayangene.com) of 1,500 exomes were further considered. Two heterozygous variants remained in \(SVEP1\) (chr9: 113137668; rs373655861; \(p.Thr3527Met\) hg19 [c.10580C\(>\)T]; GenBank: NM_153366.3) and \(UBAP1\) (chr9: 34241270; GenBank: NM_016525.4: c.436_437insTGAG \(\text{[p.Ser146Metfs*14]}\)), respectively. The \(SVEP1\) variant was ruled out by segregation studies involving Sanger sequencing of the entire pedigree. Thus, after confirmation of complete segregation, the truncating frameshift variant in ubiquitin-associated protein 1 (\(UBAP1\)) was considered as the causative allele in family 1 (Figure 1).

We then searched the GENESIS database for additional families with \(UBAP1\) variants. GENESIS contains more than 3,000 exomes and genomes from affected persons with HSP and related disorders.\(^7\) We filtered for non-synonymous and truncating variants under an autosomal-dominant model with minor-allele frequency in gnomAD < 0.0001 and a minimum sequencing depth of 10 reads. We identified seven additional HSP families, all carrying truncating variants in \(UBAP1\) (Table 1). In addition, predictively truncating \(UBAP1\) variants were prioritized in two families (9 and 10) who underwent diagnostic exome sequencing at the University of Tuebingen. The detection of truncating alleles in all families is especially remarkable when one considers the almost complete constraint of \(UBAP1\) for loss-of-function (truncating) variation in gnomAD.\(^8\)

We calculated the probability of significant enrichment of truncating variations in \(UBAP1\) in our HSP dataset compared to ExAC. In the GENESIS dataset we found seven such variants in a cohort of 567 HSP samples versus 0 truncating variants in 60,000 ExAC samples \((p = 6.187 \times 10^{-15} \text{ by Fisher test. Odds ratio = infinity)}\). Five truncating variants were reported in the \(\sim 246,000\) chromosomes in gnomAD, but none fell within the specific gene region containing the variants reported in this study.

**Figure 1.** Pedigrees of HSP-Affected Families with UBAP1 Truncations

All pedigrees suggest an autosomal-dominant or a de novo Mendelian trait. HSP-affected individuals are marked by filled symbols; individuals with unclear affection status are marked by a question mark. “mut” depicts the presence of a causative allele. Sanger traces exemplify the confirmation of variants detected via next-generation sequencing. The penetrance of truncating \(UBAP1\) variants is reduced: individual F5-III.2 was subjectively unaffected at age 14 but showed brisk reflexes of lower limbs, indicating potential dysfunction of the corticospinal tract. The 80-year-old grandfather of the index case in family 9 (F9-III.1) was unfortunately not available for a neurological examination but was reported to be in good health and without any indication of a gait disturbance.
Table 1. Detailed Genomic Locations of Detected Pathogenic Variants

<table>
<thead>
<tr>
<th>Family ID</th>
<th>Genome Assembly (hg19)</th>
<th>Isoform 1 (GenBank: NM_016525.4) Expressed in Neurons</th>
<th>Isoform 4 (GenBank: NM_001171201.1) Canonical According to NCBI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>cDNA</td>
<td>Protein</td>
</tr>
<tr>
<td>4</td>
<td>chr9: 34241384–34241384</td>
<td>c.361dupC</td>
<td>p.Leu121Profs*18</td>
</tr>
<tr>
<td>5</td>
<td>chr9: 34241396–34241396</td>
<td>c.373C &gt; T</td>
<td>p.Gln125*</td>
</tr>
<tr>
<td>6</td>
<td>chr9: 34241396–34241396</td>
<td>c.373C &gt; T</td>
<td>p.Gln125*</td>
</tr>
<tr>
<td>9</td>
<td>chr9: 34249784–34249784</td>
<td>c.1091delC</td>
<td>p.Pro364Leufs*50</td>
</tr>
</tbody>
</table>

We refer to isoform 1 throughout the text.

All additional variants and their segregation with disease in the additional families were confirmed by Sanger sequencing. On the basis of transcript GenBank: NM_016525.4, the identified variants were as follows (Table 1 and Figure 1): families 2 and 10 from Germany, c.426_427delGA (p.Lys143Serfs*15); family 3 from Canada, c.382del (p.Ser128Alafs*23); family 4 from Spain, c.361dupC (p.Leu121Profs*18); families 5 and 6 from Bulgaria (Roma ethnicity), c.373C > T (p.Gln125*); family 7 from Germany, c.286_290dupCCAGA (p.Glu97Aspfs*8); family 8 from the United States, c.295dupG (p.Asp99Glyfs*2), and family 9 from Germany c.1091delC (p.Pro364Leufs*50).

In families 2 and 4, a de novo occurrence of the truncating variant was confirmed (Figure 1). Families 5 and 6 were of self-declared Bulgarian Roma ethnicity and carried the same p.Gln125* variant, although the two index participants are from reportedly unrelated families. Evaluating the prevalence of this allele in the European Roma population and in Gypsy HSP-affected persons will require further studies.

The first manifesting symptom in all 30 UBAP1 mutation carriers from 10 families for whom detailed clinical data were available was a progressive spastic-gait disorder with a median age at onset of 8 years (interquartile range 4–9 years; oldest onset age 26 years; one asymptomatic mutation carrier (F5-III.2) aged 14 years (detailed clinical information in Table S1). At the time of examination (median disease duration 28 years; interquartile range 15–36 years), lower-limb spastic paraparesis was still the most prominent clinical feature in all affected mutation carriers; this was accompanied by brisk lower-limb tendon reflexes (all carriers, including asymptomatic carrier F5-III.2) and extensor plantar response in all but the youngest affected individual (F7-IV.6). Although brisk tendon reflexes of the upper limbs were frequently present (26 of 30; 87%) significant upper-limb spasticity was seen only in a single case (1/30; 3%; F4-II.1), consistent with a length-dependent axonopathy of the corticospinal tract. Urinary urgency was reported in some cases (11 of 30; 37%), sensory deficits were absent or mild, and there was no evidence of peripheral neuropathy. In the majority of families (8/10; 80%), no additional signs or symptoms indicating affection of neuronal systems other than the corticospinal tract were seen, and the disease was accordingly classified as pure HSP. In family 7, however, seven out of nine family members had features of cerebellar involvement (such features included saccadic pursuit, gaze-evoked nystagmus, dysmetric saccades, and limb ataxia), features also present in family 9 (F9-II.1), indicating that the cerebellum is vulnerable to UBAP1 dysfunction at least in some cases.

Overall, truncating UBAP1 mutations are associated with a predominantly pure early-onset HSP phenotype; cerebellar involvement seems to be clustered in families and was observed in 2/10 families. Although there is thus minimal variation in terms of system involvement across families carrying UBAP1 mutations, phenotypic variability exists regarding the progression rate; for example, the disease progressed rather rapidly and led to early wheelchair dependency in families 2, 6, and 7 while was almost non-progressive in family 9 (F9-II.1 is still able to run and walk unlimited distances after 38 years of disease duration). Both intrafamilial as well as interfamilial variability are common or even the norm in HSP. A complete understanding of the phenotypic spectrum associated with UBAP1 mutations will require careful clinical evaluation of additional families carrying UBAP1 mutations.

UBAP1 is a member of the endosomal sorting complex required for transport 1 (ESCRT-I) complex and a regulator of vesicular trafficking processes, binds to ubiquitinated cargo proteins, and is essential for sorting endocytic ubiquitinated cargos into multivesicular bodies (MVBs). It also plays an important role in proteasomal degradation of ubiquitinated cell-surface proteins, including EGFR (epidermal growth factor receptor) and BST2 (bone marrow...
stromal cell antigen 2). UBAP1 has two main domains: The UMA (UBAP1-MVB12-associated) domain in the N-terminal region (17–63 aa), which mediates the association with the ESCRT-I complex, and a SOUBA (solenoid of overlapping ubiquitin-associated domains) domain in the C-terminal region (389–498 aa). Both domains allow UBAP1 to act as a molecular bridge connecting the endosomal trafficking pathways to the ubiquitination machinery.

In an effort to decipher the pathophysiology of UBAP1 in HSP, we noted that all but one of the identified changes fall within a circumscribed area of the protein between Asp 99 and Ser 146; the change in family 9 at Pro364 was the only outlier (GenBank: NM_016525.4). Interestingly, disease progression in this family has been dramatically slower than in the other families: the disease has been almost stationary over decades (see above), pointing toward a possible genotype-phenotype correlation. Yet, all changes preserve the UMA domain but cause a loss of the SOUBA domain. It has been shown that mutagenesis of the SOUBA domain in UBAP1 strongly reduces its interaction with ubiquitinated proteins (Figure 2). To determine whether the observed truncating variants would lead to nonsense-mediated mRNA decay and haploinsufficiency, we evaluated both the RNA and protein expression of mutant alleles. RT-PCR was performed on RNA extracted from the fibroblasts of an affected individual, and the RNA was sequenced by the Sanger method. Surprisingly, the c.436_437insTGAG was detected in the affected person’s cDNA, indicating escape of nonsense-mediated mRNA decay (Figure S1). Next, we performed immunoblot analysis to evaluate both wild-type and potential truncated mutant UBAP1. Total protein extracts were probed with an antibody raised against the N-terminal region of UBAP1 (amino acids 25–75), a part of the protein preserved in mutant UBAP1 proteins. The protein levels measured in affected individuals were compared with those in four control fibroblasts and normalized to GAPDH levels. Immunoblots showed decreased protein levels of full-length UBAP1 in fibroblasts from affected individuals but not in control fibroblasts.
concentrations of the full-length protein in fibroblasts from affected individuals compared to controls along with the presence of the truncated protein could potentially lead to haploinsufficiency and/or a dominant-negative effect. To evaluate the effects of the truncated protein, we performed site-direct mutagenesis and generated a plasmid encoding the truncated protein fused to an HA tag at the N-terminal region. U2OS cells were co-transfected with either wild-type (HA-WT-UBAP1) or a truncated mutant (HA-Fs-UBAP1; p.Leu121Profs*18) together with its known binding partner VPS28-Myc. Both the wild-type and truncated mutant co-localize with VPS28-Myc (Figure 3A). This suggests that interaction with the ESCRT-I complex is preserved; however, the lack of the SOUBA domain, essential for ubiquitin binding, would be detrimental. Interestingly, overexpression of truncated protein containing the UMA domain has been shown to result in a dominant-negative effect by inhibiting HIV-1 budding. It is thus possible that expression of the truncated protein in affected persons could cause a dominant-negative effect due to arrest of the ESCRT-complex without acquiring the ubiquitinated protein cargo.

We performed a co-immunoprecipitation (co-IP) assay to confirm the interaction between HA-Fs-UBAP1 and VPS28-Myc. HEK293T cells were co-transfected with VPS28-Myc and with either HA-WT-UBAP1 or HA-Fs-UBAP1 and immunoprecipitated with an anti-HA or an anti-Myc antibody and analyzed by immunoblot. Our results show that both wild-type and truncated UBAP1 co-immunoprecipitated with VPS28, confirming protein-protein interaction (Figure 3B). However, ubiquitinated proteins were co-immunoprecipitated with the HA-WT-UBAP1 but not with HA-Fs-UBAP1. The arrow points to VPS28-Myc, and the asterisk below the arrow indicates the IgG band.
with HA-Fs-UBAP1. It has previously been shown that siRNA depletion of UBAP1 in HeLa cells causes clustering of early-endosome accumulation of ubiquitinated proteins and enlargement and clustering of LAMP1-positive late endosomes and lysosomes. In fibroblasts of affected persons carrying UBAP1 mutations, however, none of these changes could be observed (Figure S2), even after exposure of cells to stress conditions. It therefore appears unlikely that loss of one UBAP1 allele results in the gross failure of multivesicular body sorting.

To investigate the effects of the truncated protein in vivo, we generated a zebrafish model with UBAP1 knockdown. We used a transgenic fish with fluorescently labeled motoneuron Tg(olf2::DsRed).

Embryos were injected with CRISPR Cas9 and sgRNAs against UBAP1 supplemented with human RNA rescue of either wild-type or truncated mutant UBAP1. At 48 hours post-fertilization (hpf), embryos were imaged in vivo with a confocal microscope. We observed significantly more truncated and misshaped axons in the mutant rescued embryos than in the wild-type rescued embryos (Figure 3C). Motoneuron axon lengths in the mutant rescued embryos were imaged in vivo with a confocal microscope. We observed significantly more truncated and misshaped axons in the mutant rescued embryos than in the wild-type rescued embryos (Figure 3C). Motoneuron axon lengths in the truncated mutant were significantly (p = 0.0008) shorter than those of the wild-type (Figure 3D). This result supports the pathogenic effects of the truncated protein in vivo.

In summary, we present strong genetic evidence that truncating mutations in UBAP1 cause a relatively frequent form of HSP. UBAP1 mutations were identified in a large Iranian kindred as well as in nine additional families with different ancestral backgrounds, including Bulgarian Roma, North American of European descent, German, Spanish, and Quebecois. All available affected persons in these families carried the respective mutation in UBAP1, although UBAP1 has a strong loss-of-function constraint in the 60,000 individuals studied in the ExAC dataset. In two families we were also able to show a de novo occurrence of the variants. In our dataset of 567 families affected by dominant HSP, UBAP1 accounts for 1.2% of cases.Evaluating the full allelic and clinical spectrum in this gene will require further studies. Because UBAP1 links two cellular pathways previously involved in HSP, this finding consolidates our current understanding of the pathophysiology of HSP and points to potential novel drug targets.

Supplemental Data

Supplemental Data can be found online at https://doi.org/10.1016/j.ajhg.2019.03.001.

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Declaration of Interests

The authors declare no competing interests.

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